

FSUM 10302 PATENT

## IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Application of Robert A. Holton et al.
Scrial No. 09/063,477
Filed April 20, 1998
For PROCESS FOR THE SELECTIVE DERIVATIZATION OF TAXANES
Examiner Ba K. Trinh

TO THE COMMISSIONER OF PATENTS AND TRADEMARKS

SIR:

## DECLARATION OF ZHUMING ZHANG UNDER 37 C.F.R. 1.608(b)

- I, Zhuming Zhang, declare and state as follows:
- 1. I was a post doctorate student at Florida State University in Tallahassee, Florida. At the time of the invention, I was conducting research in Dr. Robert Holton's laboratory in the area of Synthetic Organic, Biorganic, and Organometallic Chemistry.
- 2. I am an inventor of "Process For The Selective Derivatization Of Taxanes" of application no. 09/063,477.
- 3. I conducted the "Attempt to protect C(7)OH by cbz" experiment shown on laboratory notebook page 45 before May 21, 1997 (Exhibit A). I added 53 mg of dibenzyl dicarbonate to 3 mg of 10-deacetyl baccatin III in tetrahydrofuran solvent at room temperature and allowed the mixture to react overnight. A proton NMR of the final reaction mixture revealed that 10-benzoyloxy baccatin III had been produced in greater than 90% yield.
- 4. I conducted the "Attempt to protect C(10) H by (CII<sub>3</sub>CO)-O" experiment shown on laboratory notebook page 49 before May 21, 1997 (Exhibit B). I added 10 microliters of acetic anhydride to 3 mg of 10-deacetyl baccatin III in tetrahydrofuran solvent at room temperature. I added an additional 100 microliters of acetic anhydride and allowed the mixture to react overnight. A proton NMR of the final reaction mixture revealed that baccatin III had been produced in greater than 80% yield.
- 5. I conducted the "Generation of baccatin III from 10 DAB" experiment shown on laboratory notebook page 67 before May 21, 1997 (Exhibit C). I added 1 ml of acetic anhydride to 14 mg of 10-deacetyl baccatin III and 3.5 mg ZnCl<sub>2</sub> in tetrahydrofuran solvent at room

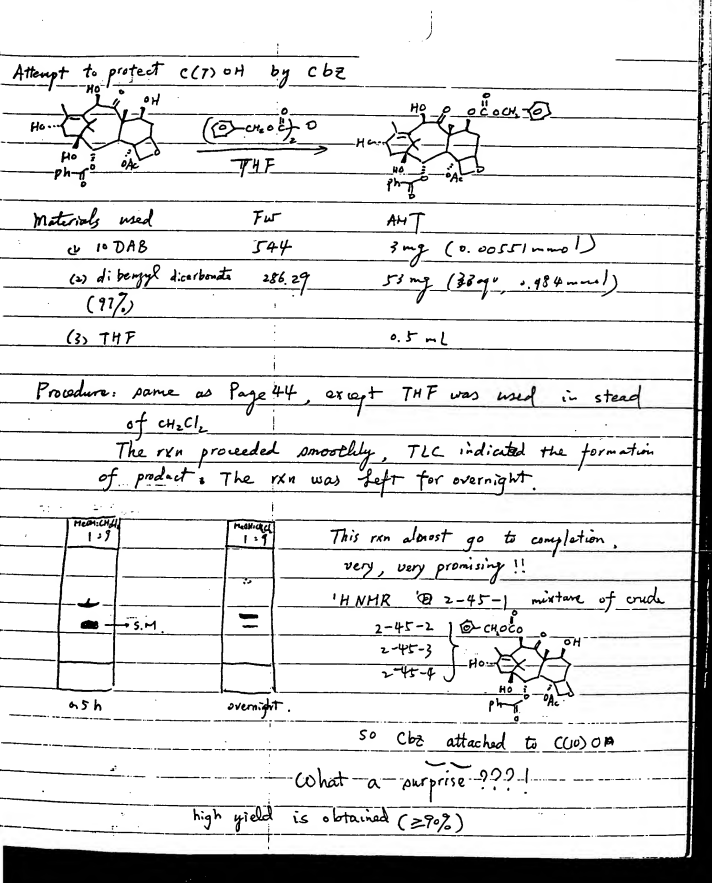
Thuming Thang
Zhuming Zhang

temperature. A proton NMR of the final reaction mixture revealed that baccatin III was a major product of the reaction.

- 6. I authored the experimental descriptions found on laboratory notebook pages 45, 49 and 67 before May 21, 1997.
- 7. I further declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further, that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under 18 U.S.C. 1001, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Date

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	- 0					
Attempt to protect ((10) H by	(c4, E) -					
04	ALO O					
Ho. Ac. O	>=\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\					
HO : JA D THE						
ph 1	ph p OAc					
700 1 1 7 7	AMT					
Materials used FW	3 mg (0,00551 mmol)					
US 10 DAB 544						
(2) AC2O 102.09	· · · · · · · · · · · · · · · · · · ·					
d=1.08						
(3) THF	o.5 mL					
Procedure: the rxn of 10DAB a	with (Phchocto suggested that similar					
reaction could take plan	ce between 10 DAB + AceO					
To a solution of 1	ODAB in THF was added Aco under hitragen.					
The reaction mixture was stirred at room temperature and						
monitored by TLC. TLC indicated the slow process of the rxn and						
a faint amount of produ	ect formed. At this stage it is not clear					
bether the ven is proceeding or	not. So 100 pl more Acro was added, After 2h,					
77. C vii	sible rxn was observed, a single spot show up					
•	high Rf value. After overnight, 280% conversion and					
monicial remand	Meanickal observed. At this stage, this					
	rxn was atopped by evaporating					
	the solvent away and taken					
	MAIN L' NAIME					
	crude 'H_NMR_2-49-1 indicated					
•.5h 2h	280 B. Intin III + 10 DAB + small					
1.5h after more Aco	amount of (7-Ac 10 DAB) ( by evaporum 7					
was added	2-49-2 pure Bacatin II. the solvent?)					

Exhibit C (Redacted)

			j	
Generation of b	raccatin I	I from los	DAB	
PO OH		: Ac	<del></del>	
ho	ACLO	: >=	OH OH	
HO 3 OALS	ZnClz	→——HO···(	<	
phy	THE		in one	
Moterials used			AHT	
CU 10 DAB		544		(0.0258 mmol)
a Aco		102.09		(0.0258 MMO)
	***	d=1.082	Inl	
(3) 2nCl2		136, 28	/	
(4) THF		130, 28		0.0258 mmol , 209v.)
(4) [7]			ImL	
Procedure. To	7117	0 4		
J - N	_a   Mr	solution of	10DAB + 2	Incl: was added ALLO
	INC AD	lution was	stirred at	- room temperature and
monitored	7 120			
TLC	actually	18 mg 2n	illz (may ha	we a little bit of H2O)
CA: Hex				
8:1	EA: Hex	EA: Hex 8:1	יא אי	1R 2-67-1 crude mixture
	<u> </u>			(too dilute)
	J22	•	Heomotia 1:7	2-67-2 crude mirture
● → SH	•SM	0	SM	
			<b>a</b>	Major baccatin II + small
				amount 7HD-diautyl
0.5h	14	1.5h		+ oxetanering opend
		· · ·		product
-	·-· <del></del>		3 h	Low temperature experiment
	···-			is recommendal.
	•			